REMARKS

Applicants have carefully reviewed the Office Action dated June 30, 2003. Claims 1-9, 11-15, 21, and 24-31 were rejected and remain pending.

The Examiner continues to reject the claims under 35 U.S.C. 35 §102(e) as being anticipated by Heck, U.S. Patent No. 6,083,207. The rejection is based on the assertion of the Examiner that the Heck patent discloses a hemostasis valve that compresses a device as that device is being passed through the hemostasis valve body. Applicants continue to traverse this rejection on multiple grounds.

First, the dilator, catheter or other medical device is not compressible in the Heck hemostasis valve and compression is not required for the valve to create a seal. Thus, the use of compression of a compressible valve sleeve to prevent flow through the valve distinguishes the current claimed invention from Heck. A second issue is that the Examiner appears to be making an argument that the Heck valve inherently compresses the device (300) as it is passed through the valve body. Because compression of the device (300) is not necessarily true, Applicants assert that this is a flawed use of the doctrine of inherency. Third, several pending claims describe a valve seat, which is not disclosed in Heck. Therefore, not every element of these claims is disclosed in Heck. Finally, the compressible valve sleeve is distinct from the dilator, catheter or other medical device in Heck and would be recognized as such by one skilled in the art. Each of these arguments highlights a difference between the current invention and Heck.

As described in each independent claim, the valve in the current invention uses compression of a compressible valve sleeve in order to restrict the flow of bodily fluids through the valve. Independent claim 1 recites, "compressing said valve sleeve for restricting fluid

flow." Independent claims 3, 12, 15, 24 and 28 have similar language describing the mechanism

for restriction of fluid flow. The Examiner has asserted that Heck anticipates these claims.

In response, Applicants continue to traverse the assertion that the Heck patent discloses a

hemostasis valve that compresses the dilator (300). Such an assertion is contrary to the manner

in which a hemostasis valve normally operates. Patent language must be read as those of

ordinary skill in the art would interpret them. See M.P.E.P. §2111.01. As a general rule, "words

in patent claims are given their ordinary meaning in the usage of the field of the invention, unless

the text of the patent makes clear that a word was used with a special meaning." Toro Co. v

White Consolidated Ind., Inc., 199 F.3d 1295, 1299 (Fed Cir. 1999). It must be determined "how

a person of experience in the field of the invention would, upon reading the patent documents,

understand the words used to define the invention." Id. Applicants respectfully assert that these

legal concepts from M.P.E.P. §2111.01 and the cases cited therein were not followed, and the

Examiner is misinterpreting Heck.

A standard hemostasis valve does not create a seal by compressing a device that is being

passed through it. As evidence of this standard definition, Applicants offer descriptions from

medical texts, the description given of a hemostasis valve in the Heck patent, and the definition

of a hemostasis valve from the patents cited in Heck. All of these sources point to one

conclusion: that a standard hemostasis valve does not create a seal by compressing that which is

being passed through it. According to M.P.E.P. §2111.01 and Toro, as stated above, Heck must

be read to use this standard definition unless Heck clearly indicates an alternate definition. Heck

does not offer such an alternate definition, and must be read to be using the standard definition of

a hemostasis valve.

The standard definition of a hemostasis valve is a valve that can accommodate a device being passed through the body of the valve. Fluid flow around the device is prevented while the device is in the valve body. In addition, fluid flow through the valve is prevented after the device is removed because the valve returns to a closed position when the device is removed from the valve body. The medical device is not pinched off in the valve body to create the seal. Instead, a flexible membrane that seals around the medical device creates the seal. Because a standard hemostasis valve seals around a device that is passed through the valve, any lumen in a device that is being passed through the valve can be kept open while the device is in the valve body. The valve is also designed to allow the device to be passed through the valve with relative ease so the operator can feel the progress of the device in the patient's body and manipulation of the device is not difficult. At the same time, the proximal end of the lumen of the device can be sealed to prevent flow through the lumen of the device itself. This standard definition of a hemostasis valve is reinforced by general literature in the field of the invention, by language in the Heck patent, and by language and Figures in the patents referred to in the Heck patent.

Hemostasis valves are defined in medical literature as valves through which a device could pass, and in the process, the hemostasis valve would form a seal <u>around</u> the device that is being passed through the valve. Examples of devices that can be passed through the body of the valve are a guide catheter or a suction thrombectomy catheter. *See Alternatives to Open*, *Vascular Surgery*, p.184 (a copy of this reference is attached for the Examiner's convenience). The fact that guide catheters or suction thrombectomy catheters can be passed through a hemostasis valve shows that a hemostasis valve does not crush the device that is being passed through the valve. With such devices that have a lumen and are passed through a hemostasis valve, it is desirable to maintain the open lumen through the device. If the lumen were not kept

open, it would eliminate the usefulness of many of the devices that are passed through the valve.

For example, a guide catheter must be kept open for another device to pass through it, and a

suction thrombectomy catheter must be kept open in order to perform the suction procedure

through the lumen. The above-cited source does not mention a seal being created by

compressing a medical device that is passed through the valve. The seal that is created by a

hemostasis valve is a seal around the device that is being passed through the valve, and the valve

is not designed to compress any such device.

In addition, an important attribute of a hemostasis valve is that a device can be passed

through the valve easily, allowing the clinician to feel the progress of the medical device through

the valve and into the patient's vasculature. Therefore, the valve must be tight enough to seal

around the device that is being passed through the valve, but not so tight as to impede the

"operator's ability to safely position or manipulate the catheter." See Cardiac Catheterization:

Application to Diagnostic and Therapeutic Procedures, p. 121 (a copy of this reference is 2

attached for the Examiner's convenience). Because compressing a device would result in an

overly tight fit between the valve and the device being passed through the valve, a hemostasis

valve is not designed to compress a device that is being passed through the valve.

These sources show that a hemostasis valve is a valve that can accommodate a device

being passed through it while sealing around, and not compressing, the device.

As shown in responses to prior Office Actions, the Heck specification and the references

used in Heck reinforce this definition. There is ample language in Heck that reinforces this

standard definition. In a response dated March 25, 2003, Applicants noted the following

language in Heck:

One method of preventing, or at least limiting, the flow of blood out of a

sheath while a pacemaker lead is being introduced is for the physician to place his

thumb over the exposed end of the sheath or to squeeze or pinch the exposed end of the sheath between his thumb and forefinger. However, neither of these methods for reducing the undesired flow of blood and air through the sheath is desirable, because the opportunity for loss of blood and introduction of air is still present. In addition, the structure of these sheaths still requires the surgeon to hold onto it while it is in place in the vessel, thereby limiting the surgeon's ability to perform other medical procedures at the same time. Moreover, squeezing the exposed end of the sheath can deform or even break the sheath, making lead insertion difficult and increasing the likelihood of damage to the lead as it passes through the sheath. Further, even when holding the end of the sheath or pinching the sheath, flow of blood out of the sheath is not entirely stopped.

Column 2, lines 14-32 (emphasis added). As Applicants noted, the underlined portions of this quotation indicate that a device being passed through the valve should not be compressed or squeezed. This reinforces the accepted definition of a hemostasis valve; a hemostasis valve forms a seal around the device and does not form a seal by compressing the device. In addition, the following language was quoted from Heck:

In addition, by sloping inward toward the lip (56), the inwardly sloped portion (60) of the outside wall (58) provides space for the lips (56) to separate without excessive force being applied, as the medical device passes through the lips (56). The inwardly sloped portion (60) of the outside wall (58) preferably slopes at an angle of about 35 to about 75 degrees from the position of the upper portion (62) so that it places pressure on the lip (56) to hold it closed against the corresponding lip of the cooperating hemostasis valve section (40), even when the medical device is forced between the lips (56).

Column 6, lines 43-53 (emphasis added). Again, instead of an indication that the medical device is to be compressed, Heck has stated that the hemostasis valve is provided with a structure specifically designed to protect the medical device against compression. The first underlined portion indicates that the hemostasis valve described by Heck is provided with a structure to reduce the compression on a medical device. The second underlined portion indicates that a medical device is advanced through the valve lips (56), and does not provide any indication of compression of the medical device. Further, this description of the operation of a hemostasis valve gives additional indications that a hemostasis valve is not intended to compress the device.

Specifically, if a hemostasis valve were meant to compress the device, it would be difficult to

pass the device through the valve, as the valve would be continuously compressing the device as

it passed through the body of the valve. Forcing the device through the valve in such a manner

would not allow the operator to "feel" the device as it is navigated through the patient's body.

This would be contrary to the description of a hemostasis valve given earlier in the Cardiac

Catheterization source, where one of the purposes of a hemostasis valve is to place minimal

pressure on the device so the operator can "feel" the progress of the device.

A third passage from Heck that was previously cited was:

Each section (38, 40) of the partitioned hemostasis valve (14) is formed from a conventional hemostasis valve material, such as a pliant, resilient rubber,

such as silicon rubber, latex rubber or a foamed rubber of 20 to 60 durometer, which can be shaped to fit within the respective body sections (26, 28) of the

partitioned hemostasis valve housing (12).

Column 5, lines 53-59. A hemostasis valve "is formed from a conventional hemostasis

valve material," which is soft and can conform around any device that is passed through the

valve. This is further reinforcement that the accepted definition of a hemostasis valve is a valve

that does not compress a device that is being passed through the valve.

All of these passages from Heck further reinforce that a hemostasis valve is a valve that

can accommodate a device being passed through it while sealing around, and not compressing,

the device.

As Applicants indicated in prior communications with the Examiner, the prior art cited in

Heck also shows that a hemostasis valve does not seal by compressing a device that is passed

through the body of the valve. Heck cites several patents that describe conventional hemostasis

valves, including U.S. Patent Nos. 5,092,857 (hereinafter Patent '857) and 4,909,798 (hereinafter

Patent '798). See Heck, column 1, lines 25-29. Both of these patents describe a hemostasis

valve as sealing <u>around</u> any devices that are passed through the valve, and neither patent mentions compressing the device that is passed through the valve body. *See* column 1, lines 37-40 of the '857 patent and column 1, lines 28-32 of the '798 patent.

Also, both the '857 patent and the '798 patent reference a body of patents that further explain the operation of a conventional hemostasis valve. Specifically, U.S. Patent Nos. 4,000,739 (hereinafter Patent '739), 4,655,752 (hereinafter Patent '752), 4,436,519 (hereinafter Patent '519), 4,610,665 (hereinafter Patent '665) and 4,673,393 (hereinafter Patent '393), and German Patent No. 30 42 229 were referenced in both the '857 patent and the '798 patent. According to the '857 patent and the '798 patent, these patents show examples of different designs of conventional hemostasis valves. *See* column 1, line 41 through column 2, line 41 of Patent '857 and column 1, line 41 through column 2, line 47 of Patent '798.

In the '739 patent, the gasket (22) sealingly engages with the catheter (46) that is passed through the valve, and it can be seen from the figures that the catheter (46) is not compressed. See column 2, lines 55-57 and Figure 2. "The slit (28) of the gasket (24) opens to allow passage of the catheter." See column 3, Lines 49-50. If the valve were meant to compress the catheter, it would not open to allow passage. In addition, "the annular gasket (22) forms a seal around the exterior of the catheter (46), thereby preventing any blood loss through the entrance hole (16)." Column 3, lines 51-53. The description of the invention mentions nothing about creating the seal by compressing the device. "The gasket (24) yields easily to the catheter and does not inhibit manipulation." Column 3, lines 53-55. This reinforces the fact that the catheter is not to be compressed. The hemostasis valve described in the '739 patent also maintains an open lumen in the catheter (46) in order to inject radiopaque material through the catheter while it is being advanced into the patient. See column 3, lines 58-60. All of this shows that the hemostasis valve

is a valve that can accommodate a device being passed through it while sealing around, and not

compressing, the device.

In the '752 patent, the hemostasis valve (referred to as a cannula) essentially acts as a

sheath for an instrument that is being passed through it. See column 4, lines 59-66. Further, it is

stated that the cannula forms a seal "about an instrument" thereby preventing fluid from passing

between the seal and the instrument and escaping out the top end of the cannula. Thus, the seal

is not formed by compressing the instrument. See column 4, lines 41-58 and column 5, lines 4-5.

It is also stated that the instrument can easily pass through the cannula, which would imply that

the instrument should not be compressed. See column 2, lines 49-52. Most importantly, Figure

6 shows a device (100) being passed through the valve and the device (100) is not being

compressed.

The '519 patent has a similar drawing that shows a device being passed through the valve

body without the device being compressed. See Figure 2. Further, the specification of the '519

patent states that fluids can be passed through the device (here, a catheter). See column 3, lines

5-6. If the device were compressed, passage of fluid through the device would be restricted or

completely occluded, as the Examiner has suggested. This would then defeat the purpose of the

catheter. There are also references in the specification to the valve forming a seal around the

catheter or device, and that the valve materials are soft, indicating further that compression is not

the mode of sealing used by a hemostasis valve. See column 3, lines 58-60 and column 4, lines

7-10.

The '665 patent has four figures that show a device being passed through the valve, and

none of these figures exhibit a device that is being compressed. See Figures 1, 3, 7 and 21. The

valve opens to conform around the catheter when the catheter is placed through the valve, and

forms a seal around the catheter. See Figure 1, 3, 7 and 21, and column 5, line 53 through

column 6, line 9. In addition, the seal material is described as a soft and pliable material and the

seal is described as being formed between the outside surface of the catheter and the seal.

Column 5, line 68 through column 6, line 9.

The '393 patent has four figures that show a device being passed through a valve, and the

device is not being compressed in any of them. See Figures 1, 3, 11 and 15. Like the '665

patent, this valve opens to allow passage of the catheter, and the valve forms a seal around the

catheter between the outside surface of the catheter and the seal. See column 4, line 28 through

column 5, line 21.

Finally, German Patent 30 42 229 also has a figure in it that shows a device passing

through the valve, and the device is not being compressed. See Figure 1 (a copy of the German

patent is attached to this Response).

Again, both the '857 patent and the '798 patent cite the above-mentioned patents as

references that describe standard hemostasis valves. See column 1, line 41 through column 2,

line 41 of the '857 patent and column 1, line 41 through column 2, line 47 of the '798 patent.

None of these references mention the compression of a device that is being passed through the

valve. In fact, all of the references either show and/or imply the opposite, in that the hemostasis

valve forms a seal around the device as the device is being passed through the valve body and

does not compress the device. Thus, both the text and the drawings of the cited patents show that

the hemostasis valve does not compress a device that is being passed through the valve body.

In summary, Applicants respectfully assert that Heck does not disclose a valve that

compresses "said valve sleeve for restricting fluid flow." See language of claim 1 from the

current invention. The standard definition of a hemostasis valve is established in the medical

literature, in the Heck patent, and in the references that are cited in Heck. That standard

definition is a valve that can accommodate a device being passed through it while sealing

around, and not compressing, the device. As mentioned earlier, M.P.E.P. §2111.01 and Toro

state that the standard definition for a word should be used unless the body of the patent clearly

states otherwise. Because Heck reinforces the standard definition and does not contradict it,

M.P.E.P. §2111.01 and Toro demand that the standard definition be used. Thus, the hemostasis

valve in Heck is a standard hemostasis valve, which is defined as a valve that can accommodate

a device being passed through it while sealing around, and not compressing, the device. It is

respectfully asserted that Heck does not disclose a valve that compresses "said valve sleeve for

restricting fluid flow." Because Heck does not include all of the elements of the independent

claims, it is respectfully asserted that all of the independent claims are allowable. Because all of

the independent claims are now allowable, all of the dependent claims are also allowable for the

same reasons and because they recite further distinct elements.

The Examiner has also stated that it must be true that the hemostasis valve of Heck

compresses the dilator (300). See page 7 of the June 30, 2003 Office Action. It appears as

though the Examiner is making an inherency argument, since an inherent characteristic is a

characteristic that must be true. See M.P.E.P. §2112. "To establish inherency, the extrinsic

evidence 'must make clear that the missing descriptive matter is necessarily present in the thing

described in the reference, and it would be so recognized by persons of ordinary skill." M.P.E.P.

§2112, citing In re Robertson, 169 F.3d 743, 745, 49 USPQ2d 1949, 1950-51 (Fed. Cir. 1999).

Thus, in the case of the Heck patent, it must necessarily be true that the device (300) is

compressed in the valve body. However, the evidence presented above leads to the conclusion

that the opposite is true, that a hemostasis valve in general, and specifically the hemostasis valve

in Heck, is a valve that can accommodate a device being passed through it while sealing around,

and not compressing, the device. Thus, it is not inherent that a hemostasis valve, including the

hemostasis valve in Heck, compresses a device that is being passed through the valve body.

Because a hemostasis valve does not inherently or otherwise compress a device that is passed

through the body of the valve, and Heck uses this standard definition, Heck does not disclose a

valve that compresses "said valve sleeve for restricting fluid flow."

It is respectfully asserted that, according to the legal doctrine from In re Robertson and

M.P.E.P. §2112, compression of a compressible valve sleeve is not inherent in Heck. Thus, all

of the elements of the current independent claims, and the claims depend therefrom, are not

disclosed by Heck, and all claims should be allowable.

Applicants also respectfully traverse the Examiner's assertion that the compressible valve

sleeve is disclosed in Heck. The Examiner has stated that the dilator, catheter or other medical

device (300) is the equivalent of the compressible valve sleeve.

However, as Applicants have indicated in past Office Actions, the compressible valve

sleeve is distinct from the dilator, catheter or other medical device that is referred to in Heck.

The claim language "must be given [its] plain meaning unless applicant has provided a clear

definition in the specification." M.P.E.P. §2111.01. "Words in patent claims are given their

ordinary meaning in the usage of the field of the invention, unless the text of the patent makes

clear that a word was used with a special meaning." M.P.E.P. §2111.01, citing In re Sneed, 710

F.2d 1544, 218 USPQ 385 (Fed. Cir. 1983).

To state that a "compressible valve sleeve" in the claims of the current invention is the

equivalent of a medical device that is referred to in Heck would give the phrase "compressible

hemostasis valve" a meaning other than the meaning attributed to the phrase by one skilled in the

art. Specifically, one skilled in the art would not compress a medical device such as a dilator or

a catheter. These medical devices do not operate well if kinks or bends are put in them, because

it will affect their pushability through the vasculature and steerability through tortuous pathways.

These devices are made to exacting standards that enable them to be effectively advanced

through a body lumen, and one skilled in the art would not knowingly make these devices less

effective by compressing them. One skilled in the art would not use a catheter for the

compressible valve sleeve because the compression of the catheter would not allow for the

operator to "feel" the progress of the catheter through the body lumen. In addition, the expense

of a device such as a catheter or a dilator would prevent one of ordinary skill in the art from

compressing such a device in the current invention when a compressible valve sleeve that is not a

medical device would suffice. Finally, when a medical device with a lumen is used, it is

desirable to keep the lumen open in order to maintain the functionality of the device.

Thus, for all of the above reasons, one skilled in the art would not use a medical device

such as a catheter or dilator in place of the compressible valve sleeve. According to M.P.E.P.

§2111.01 and In re Sneed, this definition must be used unless the patent clearly states an

alternate definition. Because the current invention does not state an alternate definition, the

definition that should be given a "compressible valve sleeve" should not be a medical device

such as a dilator or a catheter. Because Heck does not disclose a compressible valve sleeve

according to this definition and the phrase "compressible valve sleeve" is used in each

independent claim, Heck does not anticipate the independent claims or the claims dependent

therefrom, and all claims should be allowable.

Finally, Applicants respectfully assert that the rejection of claims 12, 27 and 31 is also

improper for an additional reason. Specifically, a rejection based on 35 U.S.C. §102 must be

based on prior art that discloses all of the elements of the rejected claims. "A claim is

anticipated only if each and every element as set forth in the claim is found, either expressly or

inherently described, in a single prior art reference." M.P.E.P. §2131, citing Verdegaal Bros. v.

Union Oil Co., 814 F.2d 628, 631 (Fed. Cir. 1987).

Independent claim 12 states, in part, that the valve body has a "seat for mating to said

proximal valve sleeve distal region." In addition, dependent claims 27 and 31 have similar

limitations that describe a valve sleeve seat that receives the distal end of the valve sleeve. The

valve sleeve distal end abuts this valve sleeve seat (32), and thus the valve sleeve seat does not

allow the compressible valve sleeve to be inserted into the tube (202) beyond the valve sleeve

seat. See Specification at page 7, lines 8-9 and Figure 1. There is no such valve sleeve seat in

the Heck disclosure. In fact, the dashed line in Figure 2 of Heck indicates that the device (300)

can be inserted through the valve and into the tube (202). If there were a valve sleeve seat where

the device (300) is designed to match up with the valve sleeve seat, the device (300) would not

be able to extend through the valve body and into tube (202). In addition, Figure 1 of Heck

shows the device (300) inserted in the hemostasis valve nearly up to the hub at the proximal end

of the device (300). This means that the device (300) is extending through the valve and into the

tube (202). Again, this would not be possible if the valve body contained a valve sleeve seat.

Thus, Applicants respectfully assert that Heck does not disclose an element of claims 12, 27 and

31. According to M.P.E.P. §2131 and Verdegaal, this lack of disclosure of an element shows

that Heck does not anticipate these claims, and they should be allowable.

Appl. No. 09/430,050 Response dated September 29, 2003 Reply to Office Action of June 30, 2003

Reexamination and reconsideration are respectfully requested. It is respectfully submitted that all pending claims are now in condition for allowance. Issuance of a Notice of Allowance in due course is requested. If a telephone conference might be of assistance, please contact the undersigned attorney at (612) 677-9050.

Respectfully submitted,

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By their Attorney,

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